## Food and Drug Administration, HHS

- (9) Whether the device was available for evaluation, and whether the device was returned to the manufacturer, and if so, the date it was returned to the manufacturer; and
- (10) Concomitant medical products and therapy dates. (Do not list products that were used to treat the event.)
- (d) Initial reporter information (Block E) shall contain the following:
- (1) Name, address, and phone number of the reporter who initially provided information to the user facility, manufacturer, or importer;
- (2) Whether the initial reporter is a health professional;
  - (3) Occupation; and
- (4) Whether the initial reporter also sent a copy of the report to FDA, if known.
- (e) All manufacturers (Block G) shall contain the following:
- (1) Contact office name and address and device manufacturing site;
  - (2) Telephone number;
  - (3) Report sources;
- (4) Date received by manufacturer (month, day, year);
- (5) Type of report being submitted (e.g., 5-day, initial, supplemental); and
- (6) Manufacturer report number.
- (f) Device manufacturers (Block H) shall contain the following:
- (1) Type of reportable event (death, serious injury, malfunction, etc.);
- (2) Type of followup report, if applicable (e.g., correction, response to FDA request, etc.);
- (3) If the device was returned to the manufacturer and evaluated by the manufacturer, a summary of the evaluation. If no evaluation was performed, provide an explanation why no evaluation was performed:
- (4) Device manufacture date (month, day, year);
- (5) Was device labeled for single use;
- (6) Evaluation codes (including event codes, method of evaluation, result, and conclusion codes) (refer to FDA "Coding Manual for Form 3500A");
- (7) Whether remedial action was taken and type;
- (8) Whether use of device was initial, reuse, or unknown;
- (9) Whether remedial action was reported as a removal or correction under section 519(f) of the act (list the correction/removal report number); and

- (10) Additional manufacturer narrative: and/or
  - (11) Corrected data, including:
- (i) Any information missing on the user facility report or importer report, including missing event codes, or information corrected on such forms after manufacturer verification;
- (ii) For each event code provided by the user facility under \$803.32(e)(10) or the importer under \$803.42(e)(10), a statement of whether the type of event represented by the code is addressed in the device labeling; and
- (iii) If any required information was not provided, an explanation of why such information was not provided and the steps taken to obtain such information.

[60 FR 63597, Dec. 11, 1995, as amended at 66 FR 23157, May 8, 2001]

## §803.53 Five-day reports.

A manufacturer shall submit a 5-day report to FDA, on Form 3500A or electronic equivalent as approved by FDA under §803.14 within 5 workdays of:

- (a) Becoming aware that a reportable MDR event or events, from any information, including any trend analysis, necessitates remedial action to prevent an unreasonable risk of substantial harm to the public health; or
- (b) Becoming aware of an MDR reportable event for which FDA has made a written request for the submission of a 5-day report. When such a request is made, the manufacturer shall submit, without further requests, a 5-day report for all subsequent events of the same nature that involve substantially similar devices for the time period specified in the written request. The time period stated in the original written request can be extended by FDA if it is in the interest of the public health.

## § 803.55 Baseline reports.

- (a) A manufacturer shall submit a baseline report on FDA Form 3417, or electronic equivalent as approved by FDA under §803.14 for a device when the device model is first reported under §803.50.
- (b) Each baseline report shall be updated annually, on the anniversary month of the initial submission, after the initial baseline report is submitted.